The Effects of Modified Atmosphere Packaging on the Microbiological Properties of Fresh Common Carp (Cyprinus carpio L.)

Katarína Hudecová, Hana Buchtová, Iva Steinhauserová

Department of Meat Hygiene and Technology, Faculty of Veterinary Hygiene and Ecology, University of Veterinary and Pharmaceutical Sciences Brno, Czech Republic

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Abstract

The aim of this study was to compare the growth rate of total viable counts (TVC), psychrotrophic viable counts (PVC), coliform bacteria and E. coli in portions of fresh common carp (Cyprinus carpio L.) under two different modified atmosphere packaging (experimental MAP1: 70% N2/30% CO2; experimental MAP2: 80% O2/20% CO2) and air (control samples) stored at +4 ± 0.5 °C, and to determine their shelf life. The presence of pathogens (Salmonella spp. and Listeria monocytogenes) was also surveyed in this study. A total of 360 portions from 90 common carp were examined. Laboratory analyses were performed on storage day 0 (production day) and days 3, 7 and 10. As compared to air packaging, the numbers of TVC and PVC were significantly lower (p < 0.001) in both modified atmosphere packaging (MAP) on storage days 7 and 10; coliform bacteria were significantly lower only on day 7. E. coli counts in fresh carp during storage were generally low, showing levels of < 1 log cfu/g. Salmonella spp. and Listeria monocytogenes were not detected in any of the examined samples. All the strains of Listeria spp. were identified as Listeria innocua. According to TVC values and sensory changes, the shelf life of carp portions was determined as 6 days in MAP1, 8 days in MAP2 and 3 days in air.

Freshwater fish, food safety, Listeria, Salmonella, shelf life

Fresh fish muscle is a highly perishable product with a very short shelf life, due to the presence of a large amount of water (high aw), a low content of carbohydrate (neutral pH), and the presence of native autolytic and microbial proteolytic enzymes. Spoilage of fish results from changes caused by the oxidation of lipids, reactions caused by the activities of the fish’s own enzymes, and the metabolic activities of microorganisms (Sivertsvik et al. 2002). Several methods are used to extend the shelf life of fish and fish products. The storage of fish in modified atmospheres has traditionally been used for preservation in package, and in combination with refrigeration has proven to be an effective method for extending the shelf life of fresh fish and fish products (Ruiz-Capillas and Moral 2001; Pantazi et al. 2008). Modified atmosphere packaging (MAP) is the preservation technique. In this technique, the air inside the packaging is replaced by a specific gas or mixture of gases that differ from the composition of air (Cakli et al. 2006). The gaseous atmosphere changes continuously during storage because of respiration of the packed product, biochemical changes, and the slow permeation of gases through the packaging materials (Özogul and Özogul 2006). The three main commercially used gases in modified atmosphere packaging are carbon dioxide (CO2), nitrogen (N2) and oxygen (O2). CO2 is the most important gas used in MAP for fish because of its bacteriostatic and fungistatic properties (Sivertsvik et al. 2002). The bacteriostatic effect of MAP is influenced by the CO2 concentration, the initial bacterial population, the storage temperature and the product being packaged. In food presenting high moisture and/or fat amounts, such as fish, beef and poultry, the excessive absorption of CO2 may lead to a phenomenon known as “packaging collapse”. N2 is an insipid and inert gas, showing low solubility in water and lipids. It is used for displacing the oxygen from the packaging, decreasing oxidative rancidity and inhibiting the growth of aerobic microorganisms. Due to its low solubility it is used as...
a filling gas preventing possible packaging collapse caused by the accumulation of CO₂ (Soccol and Oetterer 2003). Oxygen causes oxidative rancidity in fatty fish, stimulates growth of aerobic bacteria and inhibits growth of strictly anaerobic bacteria (Arashisar et al. 2004).

Fishery products, which are of great importance for human nutrition worldwide and provide clear health benefits, can also act as a source of food-borne pathogens (Herrera et al. 2006). Food poisoning organisms in fish are often divided into two groups: those that are naturally present in the freshwater environment, referred to as indigenous bacteria, and those associated with pollution of the aquatic environment. A third group includes bacteria introduced into fish and fish products during post-harvest handling and processing (González-Rodríguez et al. 2002). The ubiquitous nature of L. monocytogenes and Listeria spp., in conjunction with the use of surface waterways for the discharge of sewage effluents, inevitably results in the presence of these organisms in a wide range of surface water, including lakes, rivers and streams (Sauders and Wiedmann 2007). Nevertheless, the paucity of reports documenting L. monocytogenes in live freshwater fish and shellfish suggests that Listeria spp. cultured from retail products are most likely from post-harvest contamination (Wesley 2007). Listeria spp. and Listeria monocytogenes have been isolated from various species of fish and fish products around the world (Jinneman et al. 2007). The prevalence of L. monocytogenes in raw fresh fish has been analysed in many studies and varies from zero to about 30% (Miettinen and Wirtanen 2005).

Enteric organisms such as Salmonella spp. can enter aquaculture systems from many sources including farm runoff and direct contamination from wild animals, livestock and feed. These bacteria, which are one of the most important causes of human gastrointestinal disease worldwide, are not recognised as part of the normal flora of temperate aquatic environments, and their presence in aquaculture products is related to rearing practices, as well as faulty hygiene practices during post-harvest handling and processing (González-Rodríguez et al. 2002).

The aim of the present study was to observe the effect of two different modified atmospheres on the microbial growth rate in fresh chilled carp during storage, with respect to determining its shelf life and surveying the presence of Salmonella spp. and Listeria monocytogenes in common carp (Cyprinus carpio L.).

Materials and Methods

Carp production and packaging

In total, 360 samples from 90 fresh common carp were examined in this study and three different atmospheres were used: a) air (control samples), b) experimental MAP1 (70% N₂/30% CO₂), c) experimental MAP2 (80% O₂/20% CO₂).

All the samples of common carp were obtained from the same processing plant and all the carp were treated in the same way. After being killed, scaled and gutted the fish were decapitated and the fishtail trimmed. Each fish was subsequently cut into four portions of approximately the same weight. All the portions of fresh fish were primarily packed in PE bags in the processing plant and chilled to +4 ± 0.5 °C. Samples intended for storage in MAP were immediately repacked in polyamide/polyethylene pouches after the transport to the laboratory (Amilen PA/PE 20/60, VF Verpackungen GmbH, Germany) with transmission rate at 23 °C: 50 cm³/m²/day for oxygen, 10 cm³/m²/day for nitrogen and 150 cm³/m²/day for carbon dioxide. The gas mixtures used for MAP1 (70% N₂/30% CO₂) and MAP2 (80% O₂/20% CO₂) were provided by a commercial company (Linde Gas a.s., Czech Republic). After packaging the samples were stored at +4 ± 0.5 °C for 10 days.

Microbiological analyses and sensory assessment

Total viable counts (TVC), psychrotrophic viable counts (PVC), coliform bacteria and Escherichia coli were investigated on day 0 (production day) and on days 3, 7 and 10 of the storage period. Microbiological analyses were performed according to the appropriate ISO standards and a sample of 10 grams of tissue was used. After decimal dilution, total viable counts (CSN ISO 4833) and psychrotrophic viable counts (CSN ISO 17412) were determined on Standard Plate Count Agar (Oxoid, Basingstoke, UK), coliform bacteria (CSN ISO 4832) on Violet Red Bile Agar (Oxoid, Basingstoke, UK) and Escherichia coli (CSN ISO 16649-2) on TBX Agar (Merck, Darmstadt, Germany).
Determination of *Salmonella* spp. and *Listeria* spp. was performed on the production day (day 0) and on day 7 of storage. Samples of 25 grams were used for each analysis. Detection of *Salmonella* spp. was performed according to CSN EN ISO 6579 with cultivation on XLD and Brilliant Green Agar (Oxoid, Basingstoke, UK). After the enrichment procedure carried out according to CN EN ISO 11290 – 1, *Listeria* spp. were determined on Chromocult® Listeria Selective Agar acc. Ottaviani and Agosti (Merck, Darmstadt, Germany) and PALCAM Agar (Oxoid, Basingstoke, UK). Colonies suspected as *Listeria* spp. were subsequently confirmed and identified by molecular methods. After DNA isolation, genus-specific identification of isolates was primarily performed using PCR assay (Bubert et al. 1992). Strains confirmed as *Listeria* spp. were subsequently subjected to species-specific identification by multiplex PCR (Huang et al. 2007).

Sensory assessment was provided by three trained laboratory assistants. The general appearance, muscle colour and odour were examined prior to the microbiological analyses.

**Statistics**

Statistical analyses of total viable counts, psychrotrophic viable counts and coliform bacteria were performed by a Multisample Median Test (Unistat v.4.1, Unistat Ltd.). The results of microbial counts are expressed as a median with median absolute deviation (MAD). Analysis of *Listeria* spp. incidence was determined by the Mann-Whitney U-test (Statistica Cz 7, StatSoft CR s.r.o.).

**Results**

In our study, the growth rate of total viable counts, psychrotrophic viable counts and coliform bacteria in portions of fresh common carp during storage at +4 ± 0.5 °C was investigated. Fig. 1 shows total viable counts in portions of common carp stored in air, MAP1 (70% N₂/30% CO₂) and MAP2 (80% O₂/20% CO₂) during 10 days of storage. The initial microbial quality of fish was good, representing numbers of TVC of 10⁴ cfu/g. An increase in the TVC values through the storage period was observed in all types of packaging, with the most remarkable changes occurring in the control samples packaged in air. After 10 days in air packaging, the numbers of TVC reached 12.50 ± 1.00 log cfu/g. In MAP1 and MAP2 the inhibition of microbial growth was more intensive during storage when compared with air packaging. The TVC values in MAP1 and MAP2 acquired after 10 days of storage were 7.50 ± 0.20 and 7.40 ± 0.70 log cfu/g, respectively. Significant differences (p < 0.001) in TVC values between MAP1 and control samples and similarly between MAP2 and control samples were determined on storage day 7 and storage day 10. Nevertheless, no significant differences (p > 0.05) were noticed between MAP1 and MAP2 on storage days 7 and 10.

Fig. 2 shows psychrotrophic viable counts in carp portions during storage. From the initial level of 3.60 ± 0.30 log cfu/g, PVC values in air packaging increased gradually up to 11.40 ± 1.70 log cfu/g after 10 days of storage. In MAP1 the growth of psychrotrophic
microorganisms was slower, reaching figures of 6.70 ± 0.30 log cfu/g on storage day 10. PVC values showed significant differences ($p < 0.001$) between MAP1 and air packaging on storage days 7 and 10. The results of psychrotrophic viable counts in MAP2 during the ten days of storage are comparable with those of MAP1. After 10 days storage the numbers of psychrotrophic bacteria reached 7.15 ± 0.80 log cfu/g. Significant differences ($p < 0.001$) between PVC values of MAP2 and air packaging were determined on storage day 7 as well as on storage day 10.

Fig. 3 shows numbers of coliform bacteria in carp portions during storage. The contents of coliform bacteria at the beginning of storage varied in all types of packaging: 1.55 ± 0.55 log cfu/g in air, 2.10 ± 0.30 log cfu/g in MAP1 and < 1 log cfu/g in MAP2. After 7 days storage in air the numbers of coliform bacteria increased to 4.80 ± 0.25 log cfu/g, but a decrease in coliform bacteria on day 10 of storage was observed (4.15 ± 0.15 log cfu/g). Although on production day the numbers of coliforms in MAP1 were higher than in air, the growth of bacteria through the storage period was lower, showing significant differences ($p < 0.001$) on storage day 7 (3.20 ± 0.90 log cfu/g). The low numbers of coliforms detected in MAP2 on production day (< 1 log cfu/g) reached 2.30 ± 0.55 log cfu/g after 10 days of storage.
E. coli contamination in all the examined samples during this experiment was < 1 log cfu/g, except for two air-packaged samples on production day showing 1.3 and 2.3 log cfu/g respectively.

Salmonella spp. and Listeria spp. were investigated in samples on production day and after 7 days of storage (Table 1). Salmonella spp. were not detected in any of the examined samples. The number of Listeria spp. positive samples stored in air was lower on production day (15 samples) than after 7 days of storage (23 samples). In MAP1, the numbers of Listeria spp. positive samples were practically the same on storage days 0 and 7 (20 vs. 21), whereas in MAP2 the number of positive samples was lower after 7 days of storage (13 samples) than on production day (18 samples). However, no significant differences (\( p > 0.05 \)) were determined between production day and storage day 7 in any of the atmospheres. All the strains were subsequently identified as Listeria innocua.

Table 1. Incidence of Salmonella spp. and Listeria spp. in carp portions

<table>
<thead>
<tr>
<th>Type of packaging</th>
<th>No. of samples examined</th>
<th>No. of samples positive</th>
<th>Storage day 0</th>
<th>Storage day 7</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Salmonella spp.</td>
<td>Listeria spp.</td>
</tr>
<tr>
<td>Air</td>
<td>30</td>
<td>0</td>
<td>15</td>
<td>0</td>
</tr>
<tr>
<td>MAP1</td>
<td>30</td>
<td>0</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>MAP2</td>
<td>30</td>
<td>0</td>
<td>18</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>90</td>
<td>0</td>
<td>53</td>
<td>0</td>
</tr>
</tbody>
</table>

Discussion

Modified atmosphere packaging (MAP) is widely used for extending the shelf life of a wide variety of food, including fish and fish products. The efficiency of MAP in extending the shelf life of fish depends on several factors, such as the fish species, fat content, initial microbial cell load, the gas mixture in MAP and, most importantly, the storage temperature. In our study a significant decrease in the microbial growth rate was observed in MAP1 (30% CO\(_2\)/70% N\(_2\)) during storage when compared to air packaging. The effect of various CO\(_2\) contents has previously been examined in a number of studies, and delayed microbial growth has been reported e.g. in rainbow trout (Arashisar et al. 2004; Randell et al. 1997), herring (Özogul et al. 2000; Randell et al. 1997), swordfish (Pantazi et al. 2008), chub mackerel (Stamatis and Arkoudelos 2007), sea bass (Polì et al. 2006), sardines (Özogul et al. 2004), salmon (de la Hoz et al. 2000), and cod (Sivertsvik 2007). CO\(_2\) is responsible for the bacteriostatic effect in modified atmospheres and its general effects on microorganisms are the intensification of their latest growth stage and a decrease in the growth rate during the logarithmic stage (Farber 1991). However, the CO\(_2\) content in MAP used in this study was low (30%) when compared to other studies using MAP with 100% CO\(_2\) (Arashisar et al. 2004; Sivertsvik 2007), 60% CO\(_2\) (Cakli et al. 2006; Özogul et al. 2000; Özogul et al. 2004) or 50% CO\(_2\) (Ravi Sankar et al. 2008), a positive effect on reduced microbial growth was observed.

Although the application of MAP with O\(_2\) is questionable in fish because of possible lipid oxidation, a modified atmosphere with an elevated O\(_2\) level (MAP2 80% O\(_2\)/20% CO\(_2\)) was shown to be efficient in our study in terms of its inhibitory effects on the TVC and PVC. This effect on TVC and PVC was evident as early as storage day 3, although significant differences (\( p < 0.01 \)) were detected after 7 and 10 days of storage. There is little information on the effect of MAP with elevated O\(_2\) level on bacterial growth retardation in the literature. Based on microbiological, chemical and sensory analyses, Sivertsvik
Salmonella spp. are not recognised as part of the normal flora of the temperate aquatic environment and their presence in aquaculture products is related to rearing practices, as well as to faulty hygiene practices during post-harvest handling and processing (González-Rodríguez et al. 2002). In this study, Salmonella spp. were not detected in any of the examined samples. Similarly to our results, Salmonella spp. were not detected in farmed rainbow trout and salmon (González-Rodríguez et al. 2002) or in various kinds of marine fish (Davies et al. 2001; Herrera et al. 2006).

Although Listeria spp. were detected in this study, no L. monocytogenes was identified. All the strains were confirmed as L. innocua. Listeria monocytogenes has been isolated worldwide, with various prevalence ranges from a variety of fish (Ben Embarek 1994; Kwiatek 2004; Medrala et al. 2003; Miettinen and Wirtanen 2005; Herrera et al. 2006; Jinneman et al. 2007), but as in other raw foods, fishery products more frequently contain L. innocua than L. monocytogenes. Since both species share ecological niches, the presence of L. innocua is considered an indicator of possible contamination with L. monocytogenes (González-Rodríguez et al. 2002). Because of their psychrotrophic nature, Listeria spp. are able to survive and grow in fish at refrigerating temperatures even when stored under modified atmosphere packaging. Our results showed that MAP with an elevated O2 level (80%) was more effective in decreasing Listeria spp. positive samples during storage when compared to air packaging and MAP1, which can be explained by the microaerophilic and facultative anaerobic nature of Listeria spp.

No sensory changes of portions in air were detected on storage day 3. After 7 days of storage, however, changes in colour and odour were observed with an elevated amount of surface slime. These changes were even more intensive on storage day 10. In both MAP used in this study, sensory changes were only detected after 10 days of storage, accompanied by moderate colour changes, a fishy odour and a layer of surface slime.

The shelf life of carp portions was determined by total viable counts (10^6-10^7 microorganisms/g was considered the TVC limit of acceptability) and sensory changes, indicating approximately 3 days for portions in air, 6 days for MAP1, and 8 days for MAP2.
Listeria innocua. Na základe CPM a senzorických zmien bola trvanlivosť porcií kapra balených v MAP1 stanovená na 6 dní, v MAP2 na 8 dní a vo vzduchu na 3 dni.

Acknowledgement

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